Access DB# 132849

# SEARCH REQUEST FORM

## Scientific and Technical Information Center

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lf more than one search is submitted, please prioritize searches in order of need. ***********************************								
Please provide a detailed statement of the Include the elected species or structures	he search topic; and desc s, keywords, synonyms, a ns that may have a speci	cribe as specifically as possible the subject acronyms, and registry numbers, and comial meaning. Give examples or relevant of	matter to be searched.					
Title of Invention:								
Inventors (please provide full names):								
Earliest Priority Filing Date:								
*For Sequence Searches Only* Please inc. appropriate serial number.	lude all pertinent informa	ntion (parent, child, divisional, or issued paten	t numbers) along with the					
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Pate Completed: 9-21-04	Litigation		•					
earcher Prep & Review Time:		Sequence Systems						
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		Other (specify)						

PTO-1590 (8-01)



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Filtraces, review

Commend

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#### **Scientific and Technical Information Center**

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132849

Search requests relating to published applications, patent families, and litigation may be submitted by filling out this form and clicking on "Send."

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Tech Center:					
O TC 1600	0 O TC 2100 O TC 2600 O TC 2800				
O TC2900 O TC 3600	O TC 3700 C Law Lib C Other				
Enter your Contact Informat	ion below:				
Name: Jonathan Crepeau					
Employee Number: 75637	Phone: 272-1299				
Art Unit or Office: 1746	Building & Room Number: REM 6C11				
Enter the case serial number (Required): 09/939345 f not related to a patent application, please enter NA here.					
Class / Subclass(es) 429/213, 43; 361/516					
Earliest Priority Filing Date:	8/24/01				
Format preferred for results:  Paper Diskette					

### Provide detailed information on your search topic:

- In your own words, describe in detail the concepts or subjects you want us to search.
- Include synonyms, keywords, and acronyms. Define terms that have special meanings.
- \*For Chemical Structure Searches Only\*
   Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers
- \*For Sequence Searches Only\*
   Include all pertinent information (parent, child, divisional, or issued patent numbers) along with

the appropriate serial number.

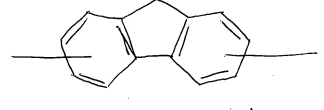
 \*For Foreign Patent Family Searches Only\* Include the country name and patent number.

Provide examples or give us relevant citations, authors, etc., if known.

 FAX or send the abstract, pertinent claims (not all of the claims), drawings, or chemical structures to your EIC or branch library.

**Enter your Search Topic Information below:** 

An electrode, preferably for a battery, capacitor, or fuel cell, comprising a polymer having the following structure contained anywhere therein:



Also note examples in the attached drawings and the language used in attached claim I ("cyclopentanone condensed w/ two aromatic rings").

**Special Instructions and Other Comments:** 

(For fastest service, let us know the best times to contact you, in case the searcher needs further clarification on your search.)

Press ALT + F, then P to print this screen for your own information.

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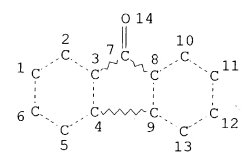
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                ACT CRE345/A
L1
                STR
L2
                SCR 2043
L3
            450 SEA FILE=REGISTRY SSS FUL L1 AND L2
     FILE 'LREGISTRY' ENTERED AT 11:53:17 ON 21 SEP 2004
L4
                STR
     FILE 'REGISTRY' ENTERED AT 12:07:22 ON 21 SEP 2004
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L6
              7 S L4 SSS FUL SUB=L3
                SAV L6 CRE345A/A
     FILE 'CAOLD' ENTERED AT 12:11:27 ON 21 SEP 2004
L7
              0 S L6
L8
              1 S L3
     FILE 'HCA' ENTERED AT 12:12:07 ON 21 SEP 2004
L9
              2 S L6
L10
            328 S L3
         726251 S ELECTROD## OR CATHOD## OR ANOD##
L11
L12
         198545 S BATTERY OR BATTERIES OR (ELECTROCHEM? OR ELECTROLY? OR
L13
         43720 S FUELCELL? OR FUEL? (2A) (CELL OR CELLS)
L14
             23 S L10 AND (L11 OR L12 OR L13)
L15
             22 S L14 NOT L9
     FILE 'REGISTRY' ENTERED AT 12:18:30 ON 21 SEP 2004
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NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 14

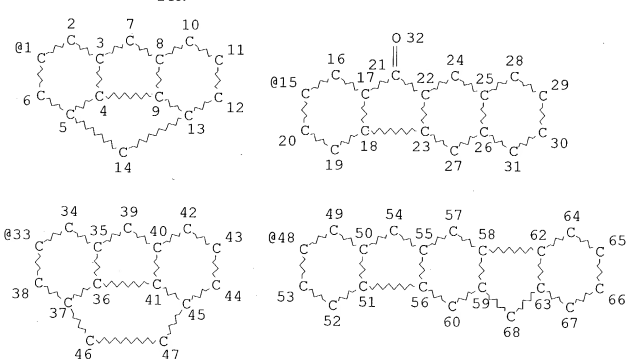
STEREO ATTRIBUTES: NONE

L2

SCR 2043

L3 450 SEA FILE=REGISTRY SSS FUL L1 AND L2

L4 STR



G1 71

VAR G1=1/15/33/48 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 68

STEREO ATTRIBUTES: NONE

L6 7 SEA FILE=REGISTRY SUB=L3 SSS FUL L4

100.0% PROCESSED 80 ITERATIONS SEARCH TIME: 00.00.01

7 ANSWERS

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FILE COVERS 1907-1966 FILE LAST UPDATED: 01 May 1997 (19970501/UP)

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L8 ANSWER 1 OF 1 CAOLD COPYRIGHT 2004 ACS on STN AN CA61:3807h CAOLD electronic interaction in ketyl radicals

=> file hca

FILE 'HCA' ENTERED AT 12:19:23 ON 21 SEP 2004
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=> d 19 1-2 ibib abs hitstr hitrn

IDISTANT KOP.

L9 ANSWER 1 OF 2 HCA COPYRIGHT 2004 ACS on STN ACCESSION NUMBER: 138:207837 HCA

TITLE:

Polymer materials for use in an electrode for use in electric energy-generating or -storing

devices

INVENTOR(S):

Umemoto, Teruo

PATENT ASSIGNEE(S):

IM & T Research, Inc., USA

SOURCE:

U.S. Pat. Appl. Publ., 21 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003044680	A1	20030306	US 2001-939345	
PRIORITY APPLN. INFO.:			US 2001-939345	200108 24
				200108

AB A carbonyl arom. polymer electrode material, suitable for use as both pos. and neg. electrodes in elec. storage devices, is disclosed. The polymers contain at least one unit having at least one cyclopentanone structure condensed with at least two arom. rings. Exemplary carbonyl arom. polymers include polymers contg. units of 9-fluorenone, cyclopenta[def]fluorene-4,8-dione, and benzo[b]fluoren-11-one. The carbonyl structure in the polymers make them very effective electrode materials which can also be anion or cation doped to increase their performance further. In addn., the polymers are proton or hydroxide anion mediators which makes them also suitable for use in electrodes in fuel cells.

IT 500149-97-3, Poly(Cyclopenta[def]fluorene-4,8-dione)

**500149-98-4**, Poly(benzo[b]fluoren-11-one)

500149-99-5, Poly(Dibenzo[b,h]fluoren-12-one)

500150-00-5, Poly(4H-Cyclopenta[def]phenanthren-4-one)

500150-02-7 500150-03-8, Poly(Indeno[1,2-

b]fluorene-6,12-dione)

(polymer materials for use in electrode for use in elec. energy-generating or -storing devices)

RN 500149-97-3 HCA

CN Cyclopenta[def]fluorene-4,8-dione, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 23702-27-4

CMF C14 H6 O2

RN 500149-98-4 HCA

CN 11H-Benzo[b]fluoren-11-one, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 3074-03-1 CMF C17 H10 O

RN 500149-99-5 HCA

CN 12H-Dibenzo[b,h]fluoren-12-one, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 53223-75-9 CMF C21 H12 O

RN 500150-00-5 HCA

CN 4H-Cyclopenta[def]phenanthren-4-one, homopolymer (9CI) (CA INDEX

NAME)

CM 1

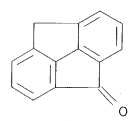
CRN 5737-13-3 CMF C15 H8 O

RN 500150-02-7 HCA

CN Cyclopenta[def]fluoren-4(8H)-one, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 500150-01-6 CMF C14 H8 O



RN 500150-03-8 HCA

CN Indeno[1,2-b]fluorene-6,12-dione, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 5695-13-6

CMF C20 H10 O2

IT 500149-97-3, Poly(Cyclopenta[def]fluorene-4,8-dione)

**500149-98-4**, Poly(benzo[b]fluoren-11-one)

**500149-99-5**, Poly(Dibenzo[b,h]fluoren-12-one)

500150-00-5, Poly(4H-Cyclopenta[def]phenanthren-4-one)

500150-02-7 500150-03-8, Poly(Indeno[1,2-

b]fluorene-6,12-dione)

(polymer materials for use in electrode for use in elec. energy-generating or -storing devices)

L9 ANSWER 2 OF 2 HCA COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

131:200157 HCA

TITLE:

Poly(indenofluorene)s (PIF) - novel conjugated

polyhydrocarbon polymers

AUTHOR(S):

Reisch, Helge A.; Schert, Ullrich

CORPORATE SOURCE:

Max-Planck-Institut fir Polymerforschung, Mainz,

D-55128, Germany

SOURCE:

Synthetic Metals (1999), 101(1-3), 128-129

CODEN: SYMEDZ; ISSN: 0379-6779

PUBLISHER:

Elsevier Science S.A.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

We are describing the synthesis of a novel class of hydrocarbon polymers, poly(indenofluorene)s PIF, composed of 3,9-di-t-butylindeno[1,2-b]fluorene- (2) or 3,7-di-t-butylindeno[2,1-b]fluorene (7) building blocks. The polymers, with high d.p. (DP: > 20), were generated by reductive coupling of the tetrachloroindenofluorene monomers 5 and 6 with low valent transition metal reagents as dehalogenating agents. PIF 2 exhibits a widely red shifted longest wavelength absorption max. (λmax: ca. 800 nm, 1,55 eV). We have also generated diketo-terminated oligomeric model compds. 9. The oligomers 9 are suitable models for an investigation of the electronic properties of the corresponding polymer PIF 2.

IT 241156-78-5P

(model oligomer; prepn. and characterization of poly(indenofluorene)s)

RN 241156-78-5 HCA

CN Indeno[1,2-b]fluoren-6(12H)-one, 12,12-dichloro-3,9-bis(1,1-dimethylethyl)-, polymer with 6,6,12,12-tetrachloro-3,9-bis(1,1-dimethylethyl)-6,12-dihydroindeno[1,2-b]fluorene (9CI) (CA INDEX NAME)

CM 1

CRN 241156-77-4 CMF C28 H26 C12 O

CM 2

CRN 150629-30-4 CMF C28 H26 C14

IT 241156-78-5P

(model oligomer; prepn. and characterization of
polv(indenofluorene)s)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d 115 1-22 cbib abs hitstr hitind

Bud date

L15 ANSWER (1) OF 22 HCA COPYRIGHT 2004 ACS on STN 141:130310 Electrochemical polymerization of 9-fluorenone. Cihaner, Atilla; Tirkes, Seha; Onal, Ahmet M. (Faculty of Engineering, Atilim University, Ankara, 06836, Turk.). Journal of Electroanalytical

Chemistry, 568(1-2), 151-156 (English) 2004. CODEN: JECHES. Publisher: Elsevier.

AB Electrochem. polymn. of 9-fluorenone, FO, was studied in CH2C12 soln. with tetrabutylammonium tetrafluoroborate, TBABF4, as the electrolyte, via const. potential electrolysis, CPE. Prior to CPE, the redox behavior of FO was studied in the same solvent/electrolytic medium using cyclic voltammetry, CV. CPE of FO yielded an insol. polymer deposit on the electrode surface. The product was characterized by FTIR spectroscopy techniques. The redox and spectroelectrochem., SPEL, behavior of the polymer were studied by CV and in situ UV-visible spectrophotometric techniques at various potentials, resp. The cond. was measured using a 2-probe technique and the paramagnetic behavior of the polymer was monitored using in situ ESR spectroscopy.

IT **500149-96-2P**, Poly(9-fluorenone)

(electrochem. prepn. and cond. and cyclic voltammetry and IR spectra of)

RN 500149-96-2 HCA

CN 9H-Fluoren-9-one, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 486-25-9 CMF C13 H8 O

CC 72-2 (Electrochemistry)

Section cross-reference(s): 35, 36

IT UV and visible spectra

(of poly(fluorenone) on ITO **electrode** in CH2Cl2 contg. tetrabutylammonium tetrafluoroborate)

IT ESR (electron spin resonance)

(of poly(fluorenone) on Pt electrode during electrochem. oxidn. and redn.)

IT Cyclic voltammetry

(of poly(fluorenone)-coated electrode in CH2Cl2 contg.
tetrabutylammonium tetrafluoroborate)

IT **500149-96-2P**, Poly(9-fluorenone)

(electrochem. prepn. and cond. and cyclic voltammetry and IR spectra of)

L15 ANSWER 2 OF 22 HCA COPYRIGHT 2004 ACS on STN

140:407358 Excitation energy transfer from polyfluorene to fluorenone defects. Gong, Xiong; Moses, Daniel; Heeger, Alan J.; Xiao, Steven (Institute for Polymers and Organic Solids, University of California at Santa Barbara, Santa Barbara, CA, 93106-9530, USA). Synthetic Metals, 141(1-2), 17-20 (English) 2004. CODEN: SYMEDZ. ISSN: 0379-6779. Publisher: Elsevier Science B.V..

Poly(9,9-dioctylfluorene-co-fluorenone) with 1% fluorenone, (PFO-F(1%)), was synthesized as a model compd. to investigate the optical and elec. effects of fluorenone defects in poly(9,9-dioctylfluorenyl-2,7-diyl), PFO. Photoluminescence (PL) studies indicate efficient Forster energy transfer from PFO to fluorenone "impurities" which are responsible for the pronounced green emission from PFO-F(1%). In electroluminescence (EL), the more pronounced green emission from PFO-F(1%) results from a combination of Forster energy transfer, charge carrier trapping, and relatively easy injection (from the electrodes) of carriers into the fluorenone traps.

IT 688318-32-3P, 2,7-Dibromo-9,9-dioctylfluorene-2,7-dibromo-9-fluorenone copolymer.

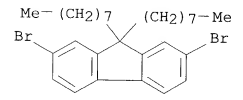
(excitation energy transfer in polydioctylfluorene contg. fluorenone moieties as model defects)

RN 688318-32-3 HCA

CN 9H-Fluoren-9-one, 2,7-dibromo-, polymer with 2,7-dibromo-9,9-dioctyl-9H-fluorene (9CI) (CA INDEX NAME)

CM 1

CRN 198964-46-4 CMF C29 H40 Br2



CM 2

CRN 14348-75-5 CMF C13 H6 Br2 O

CC 36-5 (Physical Properties of Synthetic High Polymers)
IT 688318-32-3P, 2,7-Dibromo-9,9-dioctylfluorene-2,7-dibromo-9-

fluorenone copolymer

(excitation energy transfer in polydioctylfluorene contg. fluorenone moieties as model defects)

Bud date ANSWER (3 + OF 22 L15 HCA COPYRIGHT 2004 ACS on STN Conducting polymer devices for inter-converting light and electricity. Krebs, Frederik C.; Jorgensen, Mikkel; Almdal, Kristoffer (Riso National Laboratory, Den.). PCT Int. Appl. WO 2004030029 A2 20040408, 35 pp. DESIGNATED STATES: W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG, TR. (English). PIXXD2. APPLICATION: WO 2003-EP10258 20030916. PRIORITY: GB 2002-22510 20020927.

A photovoltaic electroluminescent cell comprises a first AΒ electrode and a second electrode sepd. by dye linked block polymer mol. contg. an n-type semiconductor polymer block linked via a light absorbing monomeric dye moiety to a p-type semiconductor polymer block, the two polymer blocks being phase sepd. into distinct layers. The n-type semiconductor polymer block and the p-type semiconductor polymer block are each independently formed from selected polymers formed from terphenylenevinylene, polyaniline, polythiopene, poly(2-vinyl-pyridine), poly(N-vinylcarbazole), polyacetylene, poly(p-phenylenevinylene), polym. phenylene, poly(p-phenylene), poly(2,6-pyridine), or polypyrrole monomer, the polymers being substituted with electron withdrawing substituents in the case of the n-type polymer block and with electron donating substituents in the case of the p-type polymer block.

IT 678997-47-2P 678997-48-3P

(charge transfer complex; conducting polymer devices for inter-converting light and electricity)

RN 678997-47-2 HCA CN [1,1':4',1''-Term

[1,1':4',1''-Terphenyl]-4-acetonitrile, 4''-[2-cyano-2-(4''-formyl-2',5'-dioctyl[1,1':4',1''-terphenyl]-4-yl)ethenyl]-2',5'-dioctyl- $\alpha$ -[[4-(2,5,7-trinitro-9-oxo-9H-fluoren-4-yl)phenyl]methylene]-, compd. with  $\alpha$ -[5-[(4-mercapto-1-oxobutyl)(phenylmethyl)amino]-1-naphthalenyl]- $\omega$ -[[[[4-[[2,6-bis[3-(1-methyl-1H-phenanthro[9,10-d]imidazol-2-yl)phenyl]-4-pyridinyl]thio]phenyl](phenylmethyl)amino]carbonyl](phenylmethyl)imino]carbonyl[(phenylmethyl)imino]-1,5-naphthalenediyl] (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 678997-45-0

CMF (C25 H20 N2 O)n C91 H70 N8 O2 S2

CCI PMS

PAGE 1-A

PAGE 2-A

CM 2

CRN 677725-76-7 CMF C94 H99 N5 O8

# PAGE 1-A

Ме

#### PAGE 3-A

$$Me-(CH_2)7-Me$$

RN 678997-48-3 HCA CN [1,1':4',1''-Terphenyl]-4-acetonitrile, 4''-[2-cyano-2-(4''-formyl $2',5'-dioctyl[1,1':4',1''-terphenyl]-4-yl)ethenyl]-\alpha-[[4''-[1-cyano-2-[4-(2,5,7-trinitro-9-oxo-9H-fluoren-4-yl)phenyl]ethenyl]-2',5'-dioctyl[1,1':4',1''-terphenyl]-4-yl]methylene]-2',5'-dioctyl-, compd. with $\alpha-[5-[(4-mercapto-1-oxobutyl)(phenylmethyl)amino]-1-naphthalenyl]-\omega-[[[4-[[2,6-bis[3-(1-methyl-1H-phenanthro[9,10-d]imidazol-2-yl)phenyl]-4-pyridinyl]thio]phenyl](phenylmethyl)amino]carbonyl](phenylmethyl)imino]carbonyl[(phenylmethyl)imino]-1,5-naphthalenediyl] (1:1) (9CI) (CA INDEX NAME)$ 

CM 1

CRN 678997-45-0 CMF (C25 H20 N2 O)n C91 H70 N8 O2 S2 CCI PMS

PAGE 1-A

CM 2

CRN 677725-70-1

CMF C131 H144 N6 O8

PAGE 1-A

Ме

$$(CH_2)_7 - Me$$

$$CH$$

$$C-CN$$

$$Me$$

### PAGE 4-A

$$Me-(CH_2)7-Me$$

### IT 678997-46-1

(charge transfer complex; conducting polymer devices for

inter-converting light and electricity)

RN 678997-46-1 HCA CN [1,1':4',1''-Term

[1,1':4',1''-Terphenyl]-4-acetonitrile, 4''-formyl-2',5'-dioctyl-  $\alpha$ -[[4-(2,5,7-trinitro-9-oxo-9H-fluoren-4-yl)phenyl]methylene]-, compd. with  $\alpha$ -[5-[(4-mercapto-1-oxobutyl)(phenylmethyl)amino]-1-naphthalenyl]- $\omega$ -[[[[4-[[2,6-bis[3-(1-methyl-1H-phenanthro[9,10-d]imidazol-2-yl)phenyl]-4-pyridinyl]thio]phenyl](phenylmethyl)amino]carbonyl](phenylmethyl)imino]carbonyl[(phenylmethyl)imino]-1,5-naphthalenediyl] (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 678997-45-0

CMF (C25 H20 N2 O)n C91 H70 N8 O2 S2

CCI PMS

PAGE 1-A

PAGE 2-A

CM 2

CRN 677725-75-6 CMF C57 H54 N4 O8

PAGE 1-A

Ме

PAGE 2-A

#### ΙT 678997-44-9P

(conducting polymer devices for inter-converting light and

electricity)

RN 678997-44-9 HCA

CN Poly[(2',5'-dioctyl[1,1':4',1''-terphenyl]-4,4''-diyl)(1-cyano-1,2-ethenediyl)],  $\alpha$ -formyl- $\omega$ -[4-(2,5,7-trinitro-9-oxo-9H-fluoren-4-yl)phenyl]- (9CI) (CA INDEX NAME)

$$O_2N$$
 $NO_2$ 
 $CH_2)$   $7-Me$ 
 $CH_2)$   $7-Me$ 

IC ICM HO1L

CC 52-2 (Electrochemical, Radiational, and Thermal Energy Technology) Section cross-reference(s): 38, 74, 76

IT 678997-47-2P 678997-48-3P

(charge transfer complex; conducting polymer devices for inter-converting light and electricity)

IT 678997-46-1

(charge transfer complex; conducting polymer devices for inter-converting light and electricity)

IT 677725-70-1P 678997-19-8P 678997-44-9P 678997-45-0P (conducting polymer devices for inter-converting light and electricity)

TT 7429-90-5, Aluminum, uses 7439-95-4, Magnesium, uses 7440-00-8, Neodymium, uses 7440-02-0, Nickel, uses 7440-06-4, Platinum, uses 7440-19-9, Samarium, uses 7440-20-2, Scandium, uses 7440-22-4, Silver, uses 7440-33-7, Tungsten, uses 7440-47-3, Chromium, uses 7440-53-1, Europium, uses 7440-57-5, Gold, uses 7440-65-5, Yttrium, uses 7440-70-2, Calcium, uses 50926-11-9, Ito 157682-41-2

(electrode; conducting polymer devices for inter-converting light and electricity)

L15 ANSWER 4 OF 22 HCA COPYRIGHT 2004 ACS on STN 140:77520 Preparation and spectroelectrochemical behavior of a new alternate copolymer of 3,3'-di-n-octyl-2,2'-bithiophene and fluoren-9-one. Demadrille, Renaud; Divisia-Blohorn, Bernadette;

Zagorska, Malgorzata; Quillard, Sophie; Rannou, Patrice; Travers, Jean Pierre; Pron, Adam (DRFMC, Laboratoire de Physique des Metaux Synthetiques (CEA-CNRS-Universite Grenoble, UMR5819-SPrAM), CEA-Grenoble, Grenoble, 38054, Fr.). New Journal of Chemistry, 27(10), 1479-1484 (English) 2003. CODEN: NJCHE5. ISSN: 1144-0546. Publisher: Royal Society of Chemistry.

AB The synthesis and spectroelectrochem. behavior of a new soln. processible conjugated polymer, namely poly[{5,5'-(3,3'-di-n-octyl-2,2'-bithiophene)}-alt-(2,7-fluoren-9-one)] (abbreviated as PDOBTF), are described. PDOBTF can be considered as the first member of a new family of conjugated copolymers-poly(oligothiophene-alt-fluoren-9-one)s-whose properties can be tuned by changing the length of the oligothiophene segments and their regiochem. PDOBTF can be obtained by oxidative polymn. of 2,7-bis(4-octylthien-2-yl)-fluoren-9-one or by condensation polymn. of 2,7-bis(5-bromo-4-octylthien-2-yl)fluoren-9-one using a modification of Yamamoto coupling. prepn. methods lead to a mixt. of polymeric and oligomeric species and require post-polymn. fractionating if high mol. fractions are to be obtained. Oxidative polymn. gives product of a higher mol. wt. (Mn = 41.0 kDa, Mw/Mn = 1.81 for the highest mol. wt. fraction) as compared to the one prepd. by Yamamoto condensation polymn. (Mn =13.3 kDa, Mw/Mn = 1.45 for the highest mol. wt. fraction). Electrochem. oxidn. of PDOBTF in an nonag. electrolyte (0.1 M Bu4NBF4/acetonitrile) gives rise to an anodic peak at E = 835 mV, which can be ascribed to the p-type doping of the copolymer. The extension of the potential to  $\bar{E}$  = 1500 mV results in the oxidative degrdn. of the copolymer and induces total loss of its electroactivity. UV-Vis-NIR and Raman spectroelectrochem. data are consistent with the oxidative doping. The latter technique enables the monitoring of the doping-induced changes in both structural sub-units of the copolymer: the bithiophene sub-unit and the fluoren-9-one one.

## IT 612531-89-2P 612531-90-5P 612531-91-6P

(prepn. and spectroelectrochem. behavior of new alternate polythiophene copolymer of)

RN 612531-89-2 HCA

CN 9H-Fluoren-9-one, 2,7-bis(4-octyl-2-thienyl)-, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 612531-87-0 CMF C37 H44 O S2

RN 612531-90-5 HCA

CN 9H-Fluoren-9-one, 2,7-bis(5-bromo-4-octyl-2-thienyl)-, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 612531-88-1

CMF C37 H42 Br2 O S2

Br 
$$S$$
 Br  $S$   $Me^{-(CH_2)7}$   $(CH_2)7^{-Me}$ 

RN 612531-91-6 HCA

CN Poly[(3,3'-dioctyl[2,2'-bithiophene]-5,5'-diyl)(9-oxo-9H-fluorene-2,7-diyl)] (9CI) (CA INDEX NAME)

Me- (CH<sub>2</sub>) 7

S

(CH<sub>2</sub>) 7-Me

$$n$$

CC 35-6 (Chemistry of Synthetic High Polymers)

IT 612531-89-2P 612531-90-5P 612531-91-6P

(prepn. and spectroelectrochem. behavior of new alternate polythiophene copolymer of)

L15 ANSWER 5 OF 22 HCA COPYRIGHT 2004 ACS on STN

related app.

138:277747 Method for preparing polymers containing cyclopentanone structures. Umemoto, Teruo (IM&T Research, Inc., USA). U.S. Pat. Appl. Publ. US 2003066757 A1 20030410, 18 pp., Cont.-in-part of U.S. Ser. No. 939,141. (English). CODEN: USXXCO. APPLICATION: US 2001-23365 20011214. PRIORITY: US 2001-939141 20010824.

A method to electrolytically polymerize arom. hydrocarbons and AΒ oxidize cyclopentane structures within the hydrocarbons into cyclopentanone structures is disclosed including a method to electrolyze fluorene in the presence of an ester to produce poly(9-fluorenone). A method to electrolytically oxidize polymers having cyclopentane structures to polymers having cycloppentanone structures is also disclosed including a method to electrolyze poly(fluorene) to produce poly(9-fluorenone). These methods may include performing two sep. and independent electrolysis steps to prep. higher yield cyclopentanone structures. In addn., a method to chem. oxidize polymers contg. cyclopentane structures into polymers contg. cyclopentanone structures is disclosed, including a method to oxidize poly(fluorene), with a chem. prepd. oxidizing agent, to produce poly(9-fluorenone).

500149-96-2P, 9H-Fluoren-9-one, homopolymer (formation by electrooxidn. of fluorene in prepg. polymers contg. cyclopentanone structures)

RN 500149-96-2 HCA

CN 9H-Fluoren-9-one, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 486-25-9 CMF C13 H8 O

IC ICM C25B003-00

NCL 205429000; 205414000

CC 72-7 (Electrochemistry)

Section cross-reference(s): 38

IT 500149-96-2P, 9H-Fluoren-9-one, homopolymer (formation by electrooxidn. of fluorene in prepg. polymers contg. cyclopentanone structures)

IT 7440-02-0, Nickel, uses 7440-06-4, Platinum, uses (prepg. polymers contg. cyclopentanone structures by electrooxidn. of monomers in electrolytic cell

#### with anodes from)

L15 ANSWER 6 OF 22 HCA COPYRIGHT 2004 ACS on STN

138:46874 Double-layer formation in organic light-emitting
electrochemical cells. Ouisse, T.; Stephan, O.;
Armand, M.; Lepretre, J. C. (Laboratoire de Spectrometrie Physique,
Universite Joseph Fourier Grenoble I and CNRS (UMR C5588),
Saint-Martin d'He`res, 38042, Fr.). Journal of Applied Physics,
92(5), 2795-2802 (English) 2002. CODEN: JAPIAU. ISSN: 0021-8979.
Publisher: American Institute of Physics.

The authors present a systematic anal. of the current transients AB accompanying the formation of the electrode-electrolyte double layers in org. light-emitting electrochem. By using various room-temp. molten salts, conducting polymers, and electrodes, the current I always decreases as a power law of time,  $I \propto t-n$ . The current transients are formed of various time domains, each being characterized by a power-law exponent n<1. Impedance measurements conducted from 5 Hz to 5 MHz demonstrate that these transients represent the time response of a simple combination of const. phase angle (CPA) impedances,  $\mathrm{Zn}_{\infty}(j_{\varpi})\,\mathrm{n}$ , and of the electrolyte ionic The phys. origin of the CPA impedance is attributed to the roughness of the interface between the electrodes and the electrolyte, and to the phase sepn. within the salt-polymer blend. 107207-76-1D, Poly(9-oxo-9H-fluorene-2,7-diyl), fluorene IT

polymers contg.

(double-layer formation in org. light-emitting

(double-layer formation in org. light-emitting electrochem. cells)

RN 107207-76-1 HCA

CN Poly(9-oxo-9H-fluorene-2,7-diyl) (9CI) (CA INDEX NAME)

CC 73-11 (Optical, Electron, and Mass Spectroscopy and Other Related Properties)

Section cross-reference(s): 72

ST light emitting electrochem cell electroluminescent device; tid

IT Electric impedance

Electrochemical cells
Electroluminescent devices

(double-layer formation in org. light-emitting electrochem. cells)

107207-76-1D, Poly(9-oxo-9H-fluorene-2,7-diyl), fluorene IT polymers contq. 123863-98-9 133019-09-7, Poly(9,9-dihexyl-9H-133019-09-7D, Poly(9,9-dihexyl-9H-fluorene-2,7fluorene-2,7-diyl) diyl), fluorene polymers contg. 268536-01-2 478971-53-8 478971-56-1

> (double-layer formation in org. light-emitting electrochem. cells)

and relevent ANSWER 7 OF 22 HCA COPYRIGHT 2004 ACS on STN L15 137:370794 Preparations of electroactive fluorene-based copolymers and devices made therewith. Uckert, Frank P.; Simmons, Howard E. (E. I. Du Pont de Nemours & Co., USA). PCT Int. Appl. WO 2002090415 A1 20021114, 37 pp. DESIGNATED STATES: W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG, TR. (English). CODEN: PIXXD2. APPLICATION: WO 2002-US14094 20020503. PRIORITY: US 2001-PV288314 20010503,

The title copolymers are prepd. from: (A) a fluorene-based first ABmonomer unit, with halogenated substituents preferred, and (B) a second monomer unit selected from seven types of arom. compds., in the presence of a organometallic compd. as catalyst, and are useful for coating electron injection/transport layers in electronic devices. An example was prepd. from 2,7-diiodo-9,9-bis(2ethylhexyl) fluorene (1.08 g, 1.68 mmol) and 2,5-bis(p-bromophenyl)-N-(p-hexylphenyl)pyrrole (0.3 g, 0.56 mmol) in MePh in the presence of bis(1,5-cyclooctadiene)nickel(0) (1.23 g, 4.48 mmol), 2,2'-bipyridyl (0.70 g, 4.48 mmol), and 1,5-cyclooctadiene (0.48 g, 4.48 mmol).

IT475114-77-3P

> (prepns. of electroactive fluorene-based copolymers useful for electronic devices)

RN 475114-77-3 HCA

9H-Fluoren-9-one, 2,7-dibromo-, polymer with 2,7-dibromo-9,9-bis(2-CNethylhexyl)-9H-fluorene (9CI) (CA INDEX NAME)

CM

CRN 188200-93-3 CMFC29 H40 Br2

CM2

CRN 14348-75-5 CMFC13 H6 Br2 O

IC ICM C08G061-00

ICS C08F014-18; C08F214-18

37-3 (Plastics Manufacture and Processing) CCSection cross-reference(s): 42, 76

50926-11-9, ITO IT

> (anode; fabrication of electronic devices using electroactive fluorene-based copolymers for coating hole injection/transport layers)

5720-05-8DP, 4-Methylbenzeneboronic acid, reaction product with ΙT 1,4-benzenediboronic acid bis(neopentyl glycol) cyclic ester-2,7-diiodo-9,9-bis(2-ethylhexyl)fluorene copolymer 475114-73-9P 475114-74-0P 475114-75**-**1P 475114-76-2P 475114-77-3P 475114-78-4DP, reaction product with

4-methylbenzeneboronic acid

(prepns. of electroactive fluorene-based copolymers useful for electronic devices) and date

ANSWER 8)OF 22 HCA COPYRIGHT 2004 ACS on STN

136:254349 Organic electroluminescent device. Agata, Takashi; Okuda, Daisuke; Yoneyama, Hiroto; Seki, Mieko; Mashimo, Kiyokazu; Hirose, Eiichi; Sato, Katsuhiro; Imai, Akira; Yamamoto, Yasuo; Sugisaki, Hiroshi (Fuji Xerox Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 2002075646 A2 20020315, 16 pp. (Japanese). CODEN: JKXXAF.

APPLICATION: JP 2000-259023 20000829.

GΙ

- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT \*
- The invention relates to an org. electroluminescent device comprising org. layers sandwiched between a cathode and an anode, wherein the org. layers comprises a polymer prepd. from CH2=CR1-CO-A [R1 = H or Me group; A = I-V [X = 0, C(CN)2, C(CN)COOR2, and C(COOR2)(COOR3); Y = O and -COO(CH2)nO-; R2 and R3 = alkyl and aryl groups; R4 and R5 = alkyl, aryl, halo, etc.; W = -(CH2)nO- and -Ar-(R)k-COO(CH2)nO- [Ar = arylene; R =alkylene; k = 0 or 1]; Z = alkyl, aryl, halo, etc.; n = 1-20 integer; m and q = 0-2 integers]].
- IT 403847-72-3

(org. electroluminescent device)

RN 403847-72-3 HCA

CN 9H-Fluorene-4-carboxylic acid, 9-oxo-, 2-[(1-oxo-2-propenyl)oxy]ethyl ester, polymer with 2-hydroxyethyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 190431-71-1 CMF C19 H14 O5

CM 2

CRN 868-77-9 CMF C6 H10 O3

Februat?

$$^{\mathrm{H_{2}C}}$$
 O  $^{\parallel}$   $^{\parallel}$   $^{\parallel}$  Me-C-C-O-CH<sub>2</sub>-CH<sub>2</sub>-OH

IC ICM H05B033-14

ICS C08F020-12; C09K011-06; H05B033-22

CC 73-11 (Optical, Electron, and Mass Spectroscopy and Other Related Properties)

Section cross-reference(s): 38

IT 190431-83-5 **403847-72-3** 403847-73-4 403847-75-6 (org. electroluminescent device)

L15 ANSWER 9 OF 22 HCA COPYRIGHT 2004 ACS on STN

123:183661 Functional thin film, production and application thereof. Saji, Tetsuo (Dainichiseika Color Chem., Japan). Jpn. Kokai Tokkyo Koho JP 07062594 A2 19950307 Heisei, 41 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1993-234301 19930827.

The title film, useful for a color filter, electrophotog. device, photosensor, solar cell, electroluminescence device, optical recording device, optical nonlinear device, optoelectronic device, photochromic film, electrochromic film, gas sensor and ion sensor, is prepd. by an electrochem. redn. of a surfactant contg. an arom. azo residue, dispersed in a water or water contg. solvent. The title method requires min. or zero use of binder resin.

IT 167857-45-6

(functional thin film prepd. by photochem. redn. of surfactant contg. arom. azo residue)

RN 167857-45-6 HCA

CN Poly(oxy-1,2-ethanediyl),  $\alpha$ -[6-[4-(9-oxo-9H-fluoren-2-ylazo)phenoxy]hexyl]- $\omega$ -hydroxy- (9CI) (CA INDEX NAME)

IC ICM C25D009-08

ICS C25D013-04; G01N027-12; G01N027-333; G02B005-20; G02F001-15; G02F001-15; G02F001-17; G03G005-06; G11B007-26; H01L031-04

CC 74-13 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)
Section cross-reference(s): 52, 72, 73, 76

semicondutur device grapted as conversion device (eg. solar alls) color filter electrochemistry redn azo compd; electrophotog electrochemistry redn azo compd; photosensor electrochemistry redn azo compd; solar cell electrochemistry redn azo compd; electroluminescence electrochemistry redn azo compd; optical recording electrochemistry redn azo compd; optical nonlinear electrochemistry redn azo compd; optoelectronic device electrochemistry redn azo compd; photochromism electrochemistry redn azo compd; electrochemistry redn azo compd; gas sensor electrochemistry redn azo compd; ion sensor electrochemistry redn azo compd

ΙT 156461-29-9 156461-30-2 167856-77-1 167856-78-2 167856-79-3 167856-80-6 167856-81-7 167856-82-8 167856-83-9 167856-84-0 167856-85-1 167856-86-2 167856-87-3 167856-88-4 167856-89-5 167856-90-8 167856-91-9 167856-92-0 167856-93-1 167856-94-2 167856-95-3 167856-96-4 167856-97-5 167856-98-6 167856-99-7 167857-00-3 167857-01-4 167857-02-5 167857-03-6 167857-04-7 167857-05-8 167857-06-9 167857-07-0 167857-08-1 167857-09-2 167857-10-5 167857-11-6 167857-12-7 167857-13-8 167857-14-9 167857-15-0 167857-16-1 167857-17-2 167857-18-3 167857-19-4 167857-20-7 167857-21-8 167857-22-9 167857-23-0 167857-24-1 167857-25-2 167857-26-3 167857-27-4 167857-28-5 167857-29-6 167857-30-9 167857-31-0 167857-32-1 167857-33-2 167857-34-3 167857-35-4 167857-36-5 167857-37-6 167857-38-7 167857-39-8 167857-40-1 167857-41-2 167857-42-3 167857-43-4 167857-44-5 167857-45-6 167857-46-7 167857-47-8 167857-48-9 167857-49-0 167857-50-3 167857-51-4 167857-52-5 167857-53-6 167857-54-7 167857-55-8 167857-56-9 167857-57-0 167857-58-1 167857-59-2 167857-60-5 167857-61-6 167857-62-7 167857-63-8 167857-64-9 167857-65-0 167857-66-1 167857-67-2 167857-68-3 167857-69-4 167857-70-7 167857-71-8 167857-72-9 167857-73-0 167857-74-1

(functional thin film prepd. by photochem. redn. of surfactant contg. arom. azo residue)  $% \left( \frac{1}{2}\right) =\frac{1}{2}\left( \frac{1}{2}\right) +\frac{1}{2}\left( \frac{1}{2}\right) +\frac{1}{2}$ 

- L15 ANSWER (10) OF 22 HCA COPYRIGHT 2004 ACS on STN
  119:181376 Preparation of diphenylsilylene polymers containing
  main-chain acetylene and (hetero)aromatic groups: \(\chi(2)\) nonlinear
  optical and other properties. Corriu, Robert J. P.; Douglas,
  William E.; Yang, Zhi-xin; Karakus, Yusuf; Cross, Graham H.; Bloor,
  David (Unite Mixte CNRS/Rhone Poulenc/USTL, CNRS UMR 44, Universite
  de Montpellier II Sciences et Techniques du Languedoc, Place Eugene
  Bataillon, Montpellier, 34095/5, Fr.). Journal of Organometallic
  Chemistry, 455(1-2), 69-76 (English) 1993. CODEN: JORCAI. ISSN:
  0022-328X.
- The title polymers, (C.tplbond.CSiPh2C.tplbond.CZ)n (I; Z = p-C6H4, 4,4'-biphenylyl, 9,10-anthracenediyl, fluorenediyl, 2,2'-bipyridine-6,6'diyl, pyridinediyl, 2,5-thiophenediyl, aminonitro-m-phenylene, hydroxynitro-m-phenylene,

cyanohydroxy-m-phenylene, or p-C6F4) are prepd. by reaction of SiPh2(C.tplbond.CH)2 with the appropriate arylene dihalide in the presence of (PPh3)2PdC12, CuI and PPh3, the solvent being either NEt3 or NEt3/PhMe. The av. mol. wts. of the polymers were 2600-34,000. The UV spectra have absorption max. at 250-400 nm. The I (Z = 2-(dimethylamino)-5-nitro-m-phenylene) is  $\chi(2)$  active, r33 is 0.8 pm/V following fixed **electrode** poling at 17.5 V/ $\mu$ m. The polymers do not melt below the decompn. temp., and all transitions shown in the DSC thermogram at  $\leq$ 300° were absent on repeat scans. TGA and thermal dynamic anal. of I (Z = p-C6H4) indicated decompn. commencing at 290° and continuing to .apprx.750°. The residue was composed of  $\alpha$ -SiC and amorphous C.

IT 150303-25-6P 150321-54-3P

(prepn. and characterization of)

RN 150303-25-6 HCA

CN Poly[(diphenylsilylene)-1,2-ethynediyl(9-oxo-9H-fluorene-2,7-diyl)-1,2-ethynediyl] (9CI) (CA INDEX NAME)

$$C = C$$

$$C = C$$

$$C = C - Si$$

$$Ph$$

$$Ph$$

$$Ph$$

RN 150321-54-3 HCA

CN 9H-Fluoren-9-one, 2,7-dibromo-, polymer with diethynyldiphenylsilane (9CI) (CA INDEX NAME)

CM 1

CRN 14348-75-5 CMF C13 H6 Br2 O

CM 2

CRN 1675-57-6 CMF C16 H12 Si

```
35-5 (Chemistry of Synthetic High Polymers)
CC
ΙT
     32107-89-4P
                  131151-41-2P
                                 131151-42-3P
                                                 131151-43-4P
     131151-44-5P
                   131151-45-6P
                                  131151-46-7P
                                                 131159-95-0P
    131174-86-2P
                   131174-87-3P
                                  131174-88-4P
                                                 131174-89-5P
    131174-90-8P
                   131174-91-9P
                                  131174-92-0P
                                                 131174-93-1P
    138554-91-3P
                   150303-24-5P 150303-25-6P
                                               150303-26-7P
    150303-27-8P
                   150303-28-9P
                                  150321-52-1P
                                                 150321-53-2P
    150321-54-3P
                   150321-55-4P
                                  150321-56-5P
                                                 150321-57-6P
        (prepn. and characterization of)
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ANSWER (11) OF 22 HCA COPYRIGHT 2004 ACS on STN L15 106:127769 Redox polymer films from cathodic coupling of 4,4'-dibromobenzophenone and 2,7-dibromofluorenone. Zecchin, Sandro; Schiavon, Gilberto; Tomat, Renato; Zotti, Gianni (Ist. Polarogr. Elettrochim. Prep., C.N.R., Padua, 35020, Italy). of Electroanalytical Chemistry and Interfacial Electrochemistry, 215(1-2), 377-83 (English) 1986. CODEN: JEIEBC. ISSN: 0022-0728. Expts. were made to show the feasibility of an earlier (S. et al., AB 1984) described method for obtaining polyarom. films on The electrodeposition of electroactive films of polybenzophenone and of polyfluorenone by cathodic coupling from organonickel(II) complexes of the title compds. is described. The Ni complexes were bromo(4-bromo-4'benzophenonyl)bis(triphenylphosphine)nickel and bromo(2-bromo-7fluorencyl)bis(triphenylphosphine)nickel. The polybenzophenone and polyfluorenone films were obtained by cathodic coupling.

IT 107207-76-1P

(prepn. by cathodic coupling of nickel complexes of dibromofluorenone)

RN 107207-76-1 HCA

CN Poly(9-oxo-9H-fluorene-2,7-diyl) (9CI) (CA INDEX NAME)

CC 72-2 (Electrochemistry) Section cross-reference(s): 35

STbenzophenone fluorenone polymer cathodic coupling; dibromobenzophenone cathodic coupling polybenzophenone; dibromofluorenone cathodic coupling polyfluorenone

3988-03-2, 4,4'-Dibromobenzophenone 14348-75-5, IT2,7-Dibromofluorenone

> (cathodic coupling of nickel-coordinated, redox polymer films from)

ΙT 107282-78-0 107309-04-6

(cathodic coupling of, redox polymer films from)

ΙT

(prepn. by cathodic coupling of nickel complexes of dibromobenzophenone)

ΙT 107207-76-1P

> (prepn. by cathodic coupling of nickel complexes of dibromofluorenone)

ANSWER 12 OF 22 HCA COPYRIGHT 2004 ACS on STN

106:112096 Photoconduction in polyvinyl carbazole thin films polymerized by plasma-CVD. Inoue, Masumi; Morita, Hiromasa; Takai, Yoshiaki; Mizutani, Teruyoshi; Ieda, Masayuki (Dep. Electr. Eng., Nagoya Univ., Nagoya, 464, Japan). Japanese Journal of Applied Physics, Part 1: Regular Papers, Short Notes & Review Papers, 25(10), 1495-9 (English) 1986. CODEN: JAPNDE.

A study on the radio-frequency plasma-chem. vapor deposition of an AΒ org. photoconductor, poly(vinylcarbazole) (PVK), was carried out. The chem. structure of photoconductive PVK is similar to that of conventional PVK, from IR and UV spectroscopic studies. The voltage dependence of the photocurrent is explained by the Onsager theory. The photovoltaic effect was obsd. when Al was used as an electrode. The power-conversion efficiency was 3 +

10-3% for extremely thin photoconductive PVK film (.apprx.80 nm).

ΙT 9020-74-0

(photoconduction in thin films of, contg. trinitrofluorenone)

RN9020-74-0 HCA

9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-9H-CN carbazole homopolymer (9CI) (CA INDEX NAME)

CRN 129-79-3 CMF C13 H5 N3 O7

CM 2

CRN 25067-59-8

CMF (C14 H11 N)x

CCI PMS

CM 3

CRN 1484-13-5 CMF C14 H11 N

CC 76-5 (Electric Phenomena)

Section cross-reference(s): 74

IT 9020-74-0

(photoconduction in thin films of, contg. trinitrofluorenone)

L15 ANSWER 13 OF 22 HCA COPYRIGHT 2004 ACS on STN
104:199123 Sensor for determining arsine. Pakhomov, L. G.; Fel, Ya. A.; Soborover, E. I.; Karyakin, O. V.; Tverskoi, V. A.; Gostishchev, L. N.; Kiselev, V. D.; Pravednikov, A. N.; Pashkin, I. I.; Andrievskii, A. M. (Scientific-Research Institute of Chemistry, Gorki, USSR; Moscow Institute of Fine Chemical Technology). U.S.S.R. SU 1193557 A1 19851123 From: Otkrytiya, Izobret. 1985, (43), 181. (Russian).

CODEN: URXXAF. APPLICATION: SU 1984-3696030 19840127.

AB A copolymer of vinyl ether and 2,5,7-trinitrofluorenone-4-carboxylic acid is used to form a semiconductor layer for an AsH3 sensor.

IT 101945-03-3

(semiconductive layer from, for arsine sensor)

RN 101945-03-3 HCA

CN 9H-Fluorene-4-carboxylic acid, 2,5,7-trinitro-9-oxo-, polymer with 1,1'-oxybis[ethene] (9CI) (CA INDEX NAME)

CM 1

CRN 24929-26-8 CMF C14 H5 N3 O9

CM 2

CRN 109-93-3 CMF C4 H6 O

H<sub>2</sub>C== CH- O- CH== CH<sub>2</sub>

IC ICM G01N027-02

CC 79-2 (Inorganic Analytical Chemistry) Section cross-reference(s): 36, 76

IT Electrodes

(semiconductive, for arsine sensors)

IT **101945-03-3** 

(semiconductive layer from, for arsine sensor)

L15 ANSWER (14) OF 22 HCA COPYRIGHT 2004 ACS on STN

101:73290 Polymers from 5-membered ring hetero compounds. (Matsushita Electric Industrial Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 59043060 A2 19840309 Showa, 4 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1982-153009 19820902.

AB Polymers for elec. device applications comprise a 5-membered-ring hetero compd. and an electron acceptor. Thus, a current of 1 mA/cm2

was passed through a soln. of 1 g thiophene and 0.1 g TCNQ in 100 mL  $\,$ Me2SO4 for 30 min to form a film on the anode. The film had elec. cond. 0.43  $\Omega\text{-cm}\,.$ 

ΙT 91380-45-9P

(prepn. of, by electrochem. polymn.)

91380-45-9 HCA RN

9H-Fluoren-9-one, 2,4,7-trinitro-, polymer with 2,2'-bithiophene CN (9CI) (CA INDEX NAME)

CM1

CRN 492-97-7 CMF C8 H6 S2

CM

CRN 129-79-3 CMF C13 H5 N3 O7

C08L085-00; C08G075-00; C08G079-00; C08L081-00 IC

35-7 (Chemistry of Synthetic High Polymers) CC Section cross-reference(s): 72, 76

91380-29**-**9P

ΙT 91380-41-5P 91380-42**-**6P 91380-43-7P 91380-44-8P 91380-45-9P 91380-46-0P (prepn. of, by electrochem. polymn.)

ANSWER (15) OF 22 HCA COPYRIGHT 2004 ACS on STN 98:180014 Steady-state photoconductivity of the charge transfer complex of poly(N-vinylcarbazole) and 2,4,7-trinitro-9-fluorenone. Liu, Tongming; Wan, Meixiang; Yang, Dalin; Qian, Renyuan (Inst. Chem., Acad. Sin., Beijing, Peop. Rep. China). Gaofenzi Tongxun (4), 317-20 (Chinese) 1982. CODEN: KFTTAR. ISSN: 0453-2880.

The photoconductive spectra of poly(N-vinylcarbazole)-2,4,7-trinitro-9-fluorenone charge-transfer complex (I) [9020-74-0] (1:0.5 molar ratio) obtained with sandwich and surface cells was reported. The effects of wavelength, field strength and polarity of the illuminated electrode on Lux-Ampere index  $\gamma$  were discussed. Two kinds of photocarrier generation were suggested. When  $\lambda$  was >500 nm, the photocarriers were generated by light absorption of I; when  $\lambda$  was <500 nm, the photocarriers were produced by poly(N-vinylcarbazole) [25067-59-8] and 2,4,7-trinitro-9-fluorenone [129-79-3] free mols. present in the film.

IT 9020-74-0

(photocond. of)

RN 9020-74-0 HCA

CN 9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-9H-carbazole homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 129-79-3 CMF C13 H5 N3 O7

CM 2

CRN 25067-59-8 CMF (C14 H11 N)x CCI PMS

CM 3

CRN 1484-13-5 CMF C14 H11 N

CC 35-8 (Chemistry of Synthetic High Polymers) Section cross-reference(s): 76

IT 9020-74-0 (photocond. of)

L15 ANSWER 16 OF 22 HCA COPYRIGHT 2004 ACS on STN 98:90220 Dark conductivity of the charge transfer complex, poly(N-vinylcarbazole)-2,4,7-trinitro-9-fluorenone. Part II. Qian, Renyuan; Wan, Meixiang; Ying, Xinfang; Chen, Xiangxian (Inst. Chem., Acad. Sin., Beijing, Peop. Rep. China). Kexue Tongbao (Chinese Edition), 27(17), 1049-51 (Chinese) 1982. CODEN: KHTPAT. ISSN: 0023-074X.

AB The voltage-current curves obtained with different electrodes (Au, Ag, Cu) were superimpossible, indicating that properties in the non-Ohmic region were a body effect for 2:1 poly(N-vinylcarbazole) 2,4,7-trinitro-9-fluorenone complex [ 9020-74-0] and arose from the dependence of transition on the elec. field.

IT 9020-74-0

(cond. of, in dark, mechanism of)

RN 9020-74-0 HCA

CN 9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-9H-carbazole homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 129-79-3 CMF C13 H5 N3 O7

CRN 25067-59-8 CMF (C14 H11 N)x CCI PMS

CM 3

CRN 1484-13-5 CMF C14 H11 N

CC 36-5 (Physical Properties of Synthetic High Polymers) Section cross-reference(s): 76

IT 9020-74-0

(cond. of, in dark, mechanism of)

L15 ANSWER (17) OF 22 HCA COPYRIGHT 2004 ACS on STN 94:56526 A.S. conductance of organic solid thin films. Qian, Ren-Yuan; Jin, Xiang-Feng; Zhou, Shu-Qin (Inst. Chem., Acad. Sin., Shanghai, Peop. Rep. China). Wuli Xuebao, 29(8), 992-9 (Chinese) 1980. CODEN: WLHPAR. ISSN: 0372-736X.

The frequency dependence of a.c. conductance of org. solid thin AΒ films was studied. The dependence of the conductance on the square of the frequency is an artifact due to the electrode contact resistance and the capacitance of the film. A method based on an added external resistance is proposed to evaluate this electrode contact resistance. To characterize the a.c. conductance of org. solid thin films, 3 parameters are suggested. I.e., the d.c. specific conductance  $\sigma$ d.c. at extremely low frequencies, the exponent n in the expression  $\sigma(f) \propto fn$ for the region of higher frequencies, and the value of fx at which the straight line on the  $\lg \sigma$ - $\lg f$  plot intersects the horizontal line  $\sigma = \sigma d.c.$ . For a soln. cast film of polyvinylcarbazole (PVK)-2,4,7-trinitrofluorenone (TNF) (1:0.75) charge transfer complex and an evapd. film of Cu phthalocyanine (PcCu) the following results were obtained: PVK-TNF  $(1:0.75): \sigma d.c. = 1.9 + 10-16(\Omega.cm)-1, n = 0.96, fx$ = 6.1 + 10-3 Hz at  $14^{\circ}$ ; and PcCu:  $\sigma d.c. = 1.6$ +  $10-9(\Omega.cm)-1$ , n = 1.2, fx = 1 + 105 Hz at

17°.

IT 9020-74-0

(elec. conductance of thin films of)

RN 9020-74-0 HCA

CN 9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-9H-carbazole homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 129-79-3 CMF C13 H5 N3 O7

CM 2

CRN 25067-59-8 CMF (C14 H11 N) x

CCI PMS

CM 3

CRN 1484-13-5 CMF C14 H11 N

CC 76-2 (Electric Phenomena)

IT 147-14-8 9020-74-0

(elec. conductance of thin films of)

L15 ANSWER 18 OF 22 HCA COPYRIGHT 2004 ACS on STN 93:105800 A one-dimensional ferroelectric image sensor using an organic

photoconductor. Geary, J. M. (Bell Telephone Lab., Murray Hill, NJ, 07974, USA). Ferroelectrics, 27(1-4, 1979 IEEE Int. Symp. Appl. Ferroelectr., Part I), 231-4 (English) 1980. CODEN: FEROA8. ISSN: 0015-0193.

AB A scanning 1-dimensional image sensor was fabricated by using a Gd molybdate crystal. The ease of motion of a domain wall is perturbed by the cond. variations in a polymeric photoconductive layer coated on 1 of the crystal faces. These perturbations are sensed as alterations of the voltage-current relationship of the device as detd. by an external circuit. The device was fabricated without the use of a mask, photolithog., or electrode patterning. The optical resoln. in the direction of scanning was 20-25  $\mu.$  IT 72006-43-0

(gadolinium molybdate ferroelec. image sensor with photoconductive layer of)

RN 72006-43-0 HCA

CN 9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-3,6-diiodo-9H-carbazole homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 129-79-3 CMF C13 H5 N3 O7

CM 2

CRN 50698-53-8

CMF (C14 H9 I2 N)x

CCI PMS

CM 3

CRN 50698-52-7 CMF C14 H9 I2 N

CC 76-14 (Electric Phenomena)

Section cross-reference(s): 35, 74

IT 72006-43-0

(gadolinium molybdate ferroelec. image sensor with photoconductive layer of)

L15 ANSWER (19) OF 22 HCA COPYRIGHT 2004 ACS on STN 93:98412 Photovoltaic properties of polymer films. Reucroft, P. J.; Ullal, H. (Dep. Metall. Eng. Mater. Sci., Univ. Kentucky, Lexington, KY, 40506, USA). Solar Energy Materials, 2(2), 217-28 (English) 1980. CODEN: SOEMDH. ISSN: 0165-1633.

The effect of metal **electrode** and film thickness on the photovoltaic energy conversion efficiency in (1:1) mole ratio films of poly(N-vinylcarbazole) and 2,4,7-trinitrofluorenone was investigated. Low work function metals increase the Schottky barrier height which leads to increases in the photovoltaic energy conversion efficiency. A 10-fold decrease in film thickness produces a 103-fold increase in photovoltaic energy conversion efficiency. A theor. model which assumes that the photovoltaic current is limited by Child's law predicts photovoltaic efficiencies which are in good agreement with the measured efficiencies.

IT 39613-12-2

(photoelec. solar cells contg. films of, effect of film thickness and metal **electrodes** on properties of)

RN 39613-12-2 HCA

CN 9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-9H-carbazole homopolymer (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 129-79-3 CMF C13 H5 N3 O7

CRN 25067-59-8

CMF (C14 H11 N)x

CCI **PMS** 

> CM3

CRN 1484-13-5 CMF C14 H11 N

52-2 (Electrochemical, Radiational, and Thermal Energy Technology) CCSection cross-reference(s): 37

7440-06-4, uses and miscellaneous 7440-19-9, uses and ΙT miscellaneous 7440-22-4, uses and miscellaneous 7440-47-3, uses and miscellaneous 7440-57-5, uses and miscellaneous (electrodes, solar cells with poly(vinylcarbazole) and trinitrofluorenone films and, properties of)

ΙT 39613-12-2

> (photoelec. solar cells contg. films of, effect of film thickness and metal electrodes on properties of)

ANSWER (20) OF 22 HCA COPYRIGHT 2004 ACS on STN 85:185335 Violation of the "universality" of transit-time dispersion in 1:1 TNF.PVK. Godson, S. M.; Hirsch, J. (Birbeck Coll., Univ. London, London, UK). Solid State Communications, 20(3), 285-6 (English) 1976. CODEN: SSCOA4. ISSN: 0038-1098. AB

Transit current profiles for electrons in 1:1 TNF.PVK (Luvican) (1)

generated by an electron beam and (2) injected from an **electrode** are compared. Both sets exhibit "tails" scaling with flight-time, but the scale factors differ between the 2 sets, challenging the interpretation of the tails in terms purely of statistical flight-time dispersion.

IT 39613-12-2

(transit current profiles for electrons in)

RN 39613-12-2 HCA

CN 9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-9H-carbazole homopolymer (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 129-79-3 CMF C13 H5 N3 O7

CM 2

CRN 25067-59-8

CMF (C14 H11 N)x

CCI PMS

CM 3

CRN 1484-13-5 CMF C14 H11 N

CC 76-3 (Electric Phenomena)

IT 39613-12-2

(transit current profiles for electrons in)

ANSWER 21 OF 22 HCA COPYRIGHT 2004 ACS on STN 84:158659 Theoretical and experimental photovoltaic energy conversion in an organic film system. Reucroft, P. J.; Takahashi, K.; Ullal, H. (Dep. Metall. Eng. Mater. Sci., Univ. Kentucky, Lexington, KY, USA). Journal of Applied Physics, 46(12), 5218-23 (English) 1975. CODEN: JAPIAU. ISSN: 0021-8979.

A theor. model in which charge sepn. takes place at an AB electrode energy barrier was extended to est. the photovoltaic energy conversion efficiency as a function of film thickness and energy barrier for an org. film system based on the PVK-TNF charge-transfer complex. The model predicts that the theor. efficiency is detd. by space-charge-limited conduction for films of thickness  $>0.1~\mu.$  The efficiency is proportional to the inverse cube of the film thickness in this film thickness range. case of ultrathin films (<(0.1  $\mu)$  , the efficiency is detd. by the photocarrier generation efficiency and can approach 1%. Exptl. photovoltaic energy conversion for films of thickness 5-25  $\mu$  are in agreement with the general features predicted by the model.

ΙΤ 39613-12-2

(photovoltaic effect in films of)

RN 39613-12-2 HCA

9H-Fluoren-9-one, 2,4,7-trinitro-, compd. with 9-ethenyl-9H-CNcarbazole homopolymer (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 129 - 79 - 3CMF C13 H5 N3 O7

CM 2

25067-59-8 CRN CMF (C14 H11 N)x CCI **PMS** 

CRN 1484-13-5 CMF C14 H11 N

CC 76-9 (Electric Phenomena)
Section cross-reference(s): 52

IT 39613-12-2

(photovoltaic effect in films of)

L15 ANSWER 22 OF 22 HCA COPYRIGHT 2004 ACS on STN
59:19894 Original Reference No. 59:3547c-e Electrochemical study of free ketyl radicals. Grabowski, Zbigniew R.; Kalinowski, Marek K. (Polish Acad. Sci., Warsaw). Preprints Papers Intern. Symp. Free Radicals, 5th, Uppsala 22-1-22-7 (Unavailable) 1961.

AB Polarographic and voltammetric methods were applied to study the fluorenone system aromatic ketone-ketyl radical-aromatic pinacol i

fluorenone system aromatic ketone-ketyl radical-aromatic pinacol in aq. EtOH solns. The 1st anodic process was the oxidn. of a free radical from its redn., and the 2nd an oxidn. of its dimer, fluorenopinacol (I). The rate of dissocn. of I in solns. into radicals increased with increasing pH. In solns. of pH > 11, the radical existed as an anion. The pH independence of the redn. potential of fluorenone, and the oxidn. potential of the radical in the 1-electron oxidn.-redn. system fluorenone-fluorenone ketyl radical was  $-1.00 \pm 0.02$  v. vs. satd. calomel electrode , in 0.4M NaOH, 20 vol. % EtOH, 20°. The plot of the potential of the anodic peak of I vs. pH was a straight line with the slope dE/dpH = -0.136 v./pH, and without inflection point in the range pH 7-13.5. Oxidn. of I was a slow electron-transfer process with transmission coeff.  $\beta$  = 0.43, and preceded by an acid dissocn. reaction. Resemblances to the system benzophenone-benzophenone ketyl-benzopinacol were cited, and differences discussed.

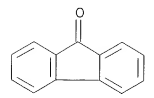
IT 101940-27-6, Fluoren-9-one, sodium ketyl, dimer (polarography of)

RN 101940-27-6 HCA

CN Fluoren-9-one, sodium ketyl, dimer (7CI) (CA INDEX NAME)

CM 1

CRN 34474-12-9 CMF C13 H8 O . Na CCI RIS



● Na+